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November 7, 2007

Robert George Kutak Rock LLP 214 W Dickson St The Three Sisters Building Fayetteville AR 72701-5221

(VIA EMAIL ONLY)

Re:

State of Oklahoma v. Tyson et al.,

No. 05-CV-0329-GKF-SAJ

#### Dear Robert:

You will find attached to this letter Bates stamped copies of the report concerning Identification of a Poultry Specific Biomarker and the November 3, 2007 report entitled Poultry-Specific Biomarker Quantitative PCR Analytical Summary. These are numbered versions of the same reports which I provided to you on Monday November 5, 2007. If you would please substitute these for the earlier production, it might help in keeping these organized.

These documents have been produced in order to assure full compliance with the Court's January 5, 2007 Order (Dkt. 1016). In providing these documents to you, we are not waiving our attorney work product privilege regarding these matters and understand the Court's Order to only relate to the production of facts and not to extend to the production of the opinions or analysis of our experts or to require further invasion of our work product privilege.

As always, should you have any questions concerning this matter, please let me know.

Sincerely,

Louis W. Bullock

## Identification of a Poultry Litter Specific Biomarker and Development of a Quantitative Assay

#### Prepared for:

Dr. Valerie J. Harwood Dept. Biology, SCA 110 University of South Florida 4202 E. Fowler Ave Tampa, FL 33620

Prepared by:



North Wind, Inc. 1425 Higham St. Idaho Falls, Idaho 83402

October 2007

NWI-3517-001

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#### **ACRONYMS**

ATCC American Type Culture Collection

DNA deoxyribonucleic acid

Case 4:05-cv-00329-GKF-PJC

MDL method detection limit

MRCF Molecular Research Core Facility

PCR polymerase chain reaction

qPCR quantitative polymerase chain reaction

T-RF terminal restriction fragment

T-RFLP terminal restriction fragment length polymorphism

UV ultraviolet

## Identification of a Poultry Litter Specific Biomarker and Development of a Quantitative Assay

#### 1 INTRODUCTION

This report documents the identification of a potential poultry litter specific biomarker, confirmation of the specificity of that biomarker for poultry litter (as compared to other fecal material), and the development of a quantitative polymerase chain reaction (qPCR) assay specific to detection of the poultry biomarker in various environmental media. Methods and results of the initial tests to identify potential poultry litter specific biomarkers are discussed in Section 2. Testing of the sensitivity of the potential biomarkers toward fecally contaminated litter and soil samples, and the specificity of the potential biomarkers against other fecal samples and various environmental media is presented in Section 3. The development and validation of an assay to quantify the poultry litter specific biomarker in various environmental media is discussed in Section 4.

### 2 IDENTIFICATION OF POTENTIAL POULTRY LITTER SPECIFIC BIOMARKER

The methods utilized to identify a potential poultry litter specific biomarker are discussed in this section. Specifically, the methods and results of microbial community profiling of poultry litter and soils to which the litter had been applied by terminal restriction fragment length polymorphism (T-RFLP) are discussed in Section 2.1. Cloning bacterial DNA from the litter and soil samples and sequencing of the plasmid deoxyribonucleic acid (DNA) containing the terminal restriction fragments (T-RFs) of interest from these clone libraries are presented in Section 2.2. Comparisons of potential biomarker DNA sequences, as compared to published sequences, and development of polymerase chain reaction (PCR) primers specific to these potential biomarkers are discussed in Section 2.3.

#### 2.1 Litter and Soil Community Profiling

#### 2.1.1 Objective

The purpose of the community profiling through the use of T-RFLP was to generate microbial community profiles of all bacteria, *Escherichia coli*, and *Bacteroides* spp. present in the poultry litter and the soils where poultry litter has been applied. The T-RFLP profiles were then used to identify common microorganisms present in both litter and soils on which poultry litter was spread.

#### 2.1.2 Methods

DNA Extraction. DNA was extracted from two poultry litter samples (five replicates each) and two agricultural soil samples (five replicates each) to which the poultry litter was applied. The poultry litter samples from which DNA was extracted were FAC-01A-1 through 5 and FAC-01B-1 through 5, and the soils samples were LAL3-A-2-1 through 5 and LAL3-B-2-1 through 5. Genomic DNA was extracted with Bio101 DNA extraction kits (QBiogene, Inc).

**T-RFLP Analysis.** T-RFLP was used to generate community profiles targeting *Bacteria* spp., *E. Coli*, and *Bacteroides* spp. The following steps were used during the T-RFLP analysis.

 The extracted DNA was PCR amplified with phosphoramidite fluorochrome 5carboxyfluorescein (FAM) labeled universal bacterial primers 8F-907R, with E.coli genus specific primers (Tsen, et al. 1998), and Bacteroides genus specific primers (Bernhard and Field, 2000). All PCR primers targeted the 16S rRNA gene.

- These PCR products were digested with the restriction enzymes Acil, Hhal, Mspl, and HaeIII (New England Biolabs).
- Common T-RFs for each PCR primer pair (e.g., universal bacteria, E.coli or Bacteroides spp.), among all 20 poultry litter and soil replicates, were targeted as potential biomarkers of poultry litter.
- T-RFLP analysis was performed by the Idaho State University, Molecular Research Core Facility (MRCF).

#### 2.1.3 Results

Common T-RFs observed in the soil and litter samples are presented in Table 1. No Bacteroides spp. fragments were identified as being present in both the soil and litter samples. T-RFs that were wellrepresented in all samples were selected for further development.

Table 1. Common T-RFs found in replicate soil and litter samples

E. coli PCR products, digested with MspI								
T-RF <sup>a</sup>	Litter FAC-01A	Litter FAC-01B	Soil LAL3-A-2	Soil LAL3-B-2				
496.0	1, <u>2</u> ,4,5 <sup>b,c</sup>	1,2,3,5	1,2,4	Present in all five				
498.9	Present in all fived	Present in all five	1,2,4,5	Present in all five				
500.8	Present in all five	Present in all five	Present in all five	Present in all five				
Universal bacteria PCR products, digested with MspI								
80.1	<u>1.2</u> ,3,4	Present in all five	Not present in any sample	1,3,4				
130.9	1,3,4	Present in all five	3	Not present in any sample				
142.9	Present in all five	1,2,3,4	1,4	1				
147.3	Present in all five	Present in all five	Present in all five	1,4				
158.9	<u>2,3</u> ,4	Present in all five	2,3,4,5	1,4				
165	1,3,4	Present in all five	1,3,4,5	1,4				

a: T-RFs of potential biomarkers are indicated in bold typeface.

b: Number indicates the litter or soil replicate sample that the T-RF was identified in.
c: An underlined number indicates that the T-RF represented <1% of community in that replicate.

d: Indicates that this T-RF was detected in all five subsamples within this sample.

#### 2.2 Cloning and Sequencing of Organisms of Interest

#### 2.2.1 Objective

The purpose of the cloning and sequencing was to obtain DNA sequences corresponding to the T-RFs of interest found in both poultry litter and soil upon which poultry litter was applied.

#### 2.2.2 Methods

- Clone libraries were constructed from the original genomic DNA extracted from the soil and litter samples and amplified with either universal bacterial or the *E. coli* genus specific primers. Four universal clone libraries were constructed from the following pooled DNA samples (i.e., 1 μl of genomic DNA extract from each sample was added to the PCR reaction for inclusion into the clones): FAC-01A-1 and FAC-01A-4, FAC-01B-3 and FAC-01B-4, LAL3-A-2-1 and LAL3-A-2-4, and LAL3-B-2-1 alone. Four *E. coli* clone libraries were constructed from the following samples: LAL3-A-2-1, FAC-01B-4, FAC-01A-4, and LAL3-B-2-1. The Topo TA cloning kit (Invitrogen) was used for construction of all clone libraries. DNA samples selected for cloning were from those subsamples in which the T-RFs of interest represented a significant portion of the fluorescence profile.
- The plasmids were excised (QIAprep Spin Miniprep Kit, QIAGEN) from the clones and analyzed by T-RFLP (digested with MspI alone) to determine which clones contained the T-RFs of interest (see Table 1).
- Plasmids containing the T-RFs of interested were amplified by PCR and sequenced using the primers T7, T3, 519R, and 338F for double coverage of the 16s rDNA.

#### 2.2.3 Results

- Only three of the six T-RFs representing potential biomarkers were found in the universal clone library (i.e., T-RFs 142.9, 147.3, and 158.9). Sequences representing all three E. coli T-RFs were found in the E. coli clone library.
- After developing two clone libraries and screening an additional 88 clones with T-RFLP, the target biomarkers with T-RFs of 80.1, 130.6 and 165 were not found. A total of 350 clones were screened.
- T-RFLP sequence analysis and DNA sequencing of each clone was performed by the MRCF.

#### 2.3 Biomarker Sequence Analysis and PCR Primer Design

#### 2.3.1 Objective

The purpose of the biomarker sequence analysis was to compare the potential biomarker DNA sequences to published 16S rRNA DNA sequences to determine whether these sequences had previously been observed, and whether they were identified as a particular species or associated with a particular host animal. Additionally, these sequences were used to design PCR primers specific to each biomarker.

#### 2.3.2 Methods

- The three universal and the three *E. coli* sequences corresponding to the T-RFs of interest were compared to the BLAST database (i.e., National Center for Biotechnology Information) to determine closely related organisms to our potential biomarkers and sites amenable for the design of PCR primers.
- PCR primers were designed for the three biomarkers from the universal bacterial library and one of the E. coli biomarkers, targeting regions of variability between our sequence and the database sequences of the top 20 matches in the BLAST database. PCR primers were designed using the Primer Express v2.0 software (Applied Biosystems). These primers were analyzed for thermodynamic folding problems and compared to the RDPII database (Michigan State University) to determine what other organisms they might amplify. The results of the analysis of the forward and reverse PCR primer comparison against the RDPII database are presented in the Table 2.

Table 2. Primer design for potential biomarkers identified during T-RFLP/clone library assessment.

Potential Biomarker Clone Number/ T-RF/Organism *	Forward Primer Accession Number of Closest Match in Sequence Database RPDII	Reverse Primer Accession Number of Closest Match in Sequence Database RPDII
Clone SA19 T-RF-158.9 Kineococcus	AY919955, AY426452, AF195447, AF513961, AY862810, AM085954, AM182287, AM182297, AM182298, AM182299	Primer sequence did not match any organisms in the database
Clone SB37 T-RF 142.9 Uncultured organisms	Primer sequence did not match any organisms in the database	Primer sequence did not match any organisms in the database
Clone LA35 T-RF 147.3 Brevibacterium	Primer sequence did not match any organisms in the database	Primer sequence did not match any organisms in the database
Clone SA15 T-RF 500.8 Pantoea ananatis	Primer sequence did not match any organisms in the database	AJ010486, DQ221344, AF364845, AF364844, AY528223, AY579209    AY579211, U80196, U80209, AB004758, AB027693, AY530796    AJ629190, AB178169, AB178170, AY898643, AB114622, DQ133548    DQ195524, AB242937, AB242945    AB242946, AB242979
		And an additional 30 sequences  Enterobacter spp.

#### 2.3.3 Results

• Only one E. coli biomarker had a variable region observed in all 20 matches of the closest related organisms in the BLAST database. PCR primers were designed for this variable

region and another region that had a mismatch between our sequence and the database sequence in 10 of the top 20 matching sequences.

- One primer pair was designed that was determined to be specific for T-RF of interest 147.3 from clone LA-35, a Brevibacterium spp., and would have no matches compared to the sequences in the RDPII database.
- One primer pair was designed for a Kineococcus spp. corresponding to T-RF 158.9, where the forward primer matched the sequences of 10 other organisms in the RDPII database, but the reverse primer was specific to this Kineococcus spp.
- One primer pair was designed for an organism not matching any cultured organisms in the BLAST or RDPII databases corresponding to T-RF 142.9.

#### 3 TESTING OF THE SENSITIVITY AND SPECIFICITY OF THE POTENTIAL POULTRY BIOMARKER TARGETS

The methods utilized to test the sensitivity and specificity of the potential poultry litter specific biomarkers are discussed in this section. Specifically, the methods and results for the testing the PCR primers in the original soil and litter samples used for the T-RFLP are presented in Section 3.1, while the testing of the PCR primers for the biomarkers against closely related organisms in the BLAST database search are presented in Section 3.2. Testing of the PCR primers for the various potential biomarkers against other fecal material is presented in Section 3.3. Cloning and sequencing of PCR amplicons derived from fecal samples that amplified with the Brevibacteria specific primers and comparison to the biomarker sequence is presented in Section 3.4 Finally, testing of the PCR primers of the confirmed poultry litter specific biomarker in environmental samples from the potentially poultry litter impacted watershed are presented in Section 3.5.

#### Test PCR Primers Against Original Soil and Litter Samples 3.1

#### 3.1.1 Objective

The objective of this test was to determine if the PCR primers specific for the various biomarkers amplified DNA from the original contaminated soil and poultry litter samples used to find the T-RFs of interest (i.e., potential biomarker sequences).

#### 3.1.2 Methods

- A nested PCR approach was used to increase sensitivity of the PCR assay by first amplifying the DNA with the universal bacterial primers 8F-907R or E. coli speciesspecific primers, purifying the PCR products (OIAquick PCR purification kits, OIAGEN). and then amplifying the DNA with the biomarker-specific primers.
- PCR products were separated by gel electrophoresis, stained with ethidium bromide, and visualized by ultraviolet (UV) light.

#### 3.1.3 Results

The results of the nested PCR with the potential biomarker-specific PCR primers of DNA from the original litter and soil samples are presented in Table 3.

. Table 3. Test of the biomarker-specific PCR analysis on original litter and soil replicates.

	Clone LA35 Clone SB37 Brevibacterium spp. Unknown genus			e SA19 occus spp.		Clone SA15 E. coli		
Sample	% of T-RF Profile	Amplified with LA35 Primers?	% of T-RF Profile	T-RF with SB37		Amplified with SA19 Primers?	% of T-RF Profile	Amplified with SA15 Primers?
			Litte	r Sample Resu	lts			
FAC-01-A-1	2.6	yes	2.8	yes	not present <sup>b</sup>	not run	26.4	yes
FAC-01-A-2	2.6	yes	4.6	no	0.8	yes	13	no
FAC-01-A-3	3.4	yes	5.4	yes	1	not run <sup>c</sup>	no data*	yes
FAC-01-A-4	3.3	yes	5.9	yes	1.3	not run <sup>c</sup>	23.3	по
FAC-01-A-5	no data*	yes	no data*	not run <sup>e</sup>	no data *	not run <sup>e</sup>	28.5	yes
FAC-01-B-1	3.3	yes	3.6	not run <sup>c</sup>	1.5	not run <sup>c</sup>	33	yes
FAC-01-B-2	4	yes	5.5	yes	1.1	no	43.8	yes
FAC-01-B-3	3.5	yes	6.8	yes	1.3	yes	16.5	yes
FAC-01-B-4	3.2	yes	5.7	not run <sup>c</sup>	1.2	not run <sup>c</sup>	29.1	no
FAC-01-B-5	4,5	yes	8	not run°	1.1	not run <sup>e</sup>	48.5	no
			Soil	Sample Result	S			
LAL3-A-2-1	7.1	yes	0.8	yes	2.3	not run°	6.7	yes
LAL3-A-2-2	12.7	no	not present <sup>b</sup>	yes	3.4	not run <sup>c</sup>	14.3	по
LAL3-A-2-3	9	yes	1	yes	3.6	not run <sup>e</sup>	25.3	yes
LAL3-A-2-4	6.9	yes	0.8	yes	3.2	yes	10.4	yes
LAL3-A-2-5	9.5	no	0.9	yes	3.6	not run <sup>e</sup>	2.5	yes
LAL3-B-2-1	6	yes	0.8	yes	3	yes	18.1	yes
LAL3-B-2-2	no dataª	yes	no dataª	yes	no data"	yes	6.8	yes
LAL3-B-2-3	not present <sup>b</sup>	yes	not present <sup>b</sup>	yes	3	no	9.1	yes
LAL3-B-2-4	6.3	yes	0.8	yes	3.5	no	2.6	yes
LAL3-B-2-5	no data	yes	no data*	yes	no data*	yes	7.1	yes

		e LA35 terium spp.		ie SB37 wn genus				
Sample_	% of T-RF Profile	Amplified with LA35 Primers?	% of T-RF Profile	Amplified with SB37 Primers?	% of T-RF Profile	Amplified with SA19 Primers?	% of T-RF Profile	Amplified with SA15 Primers?

a No data indicates that the T-RFLP analysis was not completed on this sample.

#### 3.2 Test PCR Primer Set LA35 Against a Closely Related Bacterium

#### 3.2.1 Objective

The objective of this test was to determine if the PCR primers for the Brevibacterium LA35 potential poultry litter biomarker amplified the same product in Brevibacterium sp. CHNDP32 (DQ337537), the fourth closest related organism found in the BLAST search.

#### 3.2.2 Methods

- 16S rDNA of the closely related organism identified in the BLAST search was obtained from Dr. Chee-Sanford from the University of Illinois at Urbana-Champaign.
- The Brevibacterium sp. CHNDP32 was PCR amplified using the LA35 primers, and PCR products were separated by gel electrophoresis, stained with ethidium bromide, and visualized by UV light.

#### 3.2.3 Results

The Brevibacterium biomarkers primers did not amplify the Brevibacterium sp. CHNDP32.

#### 3.3 Test PCR Primers Against Other Fecal Samples from Within and Outside the Watershed

#### 3.3.1 **Objective**

The purpose of this test was to determine the specificity of the four potential poultry biomarker targets (LA35, SB37, SA19 and SA15) against other sources of fecal contamination from within and outside the potentially poultry impacted watershed.

#### 3.3.2 Methods

- Fecal samples were collected in duplicate from beef and dairy cattle, swine, geese, ducks, and humans from inside and outside the potentially poultry impacted watershed. Field blank controls were included with each type of fecal sample. The fecal samples were preserved in glycerol and shipped on ice to the laboratory.
- Genomic DNA was extracted from all fecal samples using the Bio 101 FastDNA SPIN Kit for Soil, DNA extraction kit (Qbiogene, Inc).

b Not present indicates that the organism represented by that T-RF was not found in the original analysis (see Section 2.1). c Not run indicates that this sample was not run with PCR.

#### 3.3.3 Results

The results of the PCR amplification of the fecal samples with the four potential poultry litter biomarkers are presented in Table 4.

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		:		Did the Potential B	Did the Potential Biomarker Sequence PCR Amplify in the Fecal Sample?	PCR Amplify in th	e Fecal Sample?
				_	PARCY TITLE		
	Type of Fecal	Inside or Outside the		Clone LA35	Unknown	Clone SA19	Clone SA15
Sample	Sample	Watershed	City	Brevibacterium	Organism	Kineococcus	E. coli
MAN-BC-1-a		Outside		No	Νο	No	No
MAN-BC-1-b		Outside		No	Š	No	No
MAN-BC-2-a		Outside		SN	Yes	SN N	SZ.
MAN-BC-2-b		Outside		SN S	Yes	No	No
MAN-BC-3-a		Outside		No	No	No	No
MAN-BC-3-b		Outside		No	No No	No	Š
MAN-BC-4-a		Outside		No	%	No	Š
MAN-BC-4-b		Outside		No	No	No	Š
MAN-BC-5-a		Outside		No	Yes	Yes	S <sub>S</sub>
MAN-BC-5-b	Reef Cattle	Outside		No	N <sub>o</sub>	No	Š
MAN-BC-6-a		Inside		No	Yes, faint	No	%
MAN-BC-6-b		Inside		No	No	No	ž
MAN-BC-7-a		Inside		No	Yes	Yes, faint	Yes
MAN-BC-7-b		Inside		No a	Yes	Yes	Yes
MAN-BC-8-a		Inside		No	No	Yes	%
MAN-BC-8-b		Inside		No	No	Yes, faint <sup>b</sup>	No
MAN-BC-9-a		Inside		No	No	No	No
MAN-BC-9-b	1	Inside		No	No	Yes, faint <sup>b</sup>	No
MAN-BC-10-a		Inside		No	Yes, faint <sup>b</sup>	Yes, faint	No
MAN-BC-10-b		Inside		No	Yes	Yes	No
MAN-BC-F-a	Field Blank	Outside		No	Yes	No	No
MAN-DC-1		1					
1200		Ourside		8	Yes	Yes, faint	Yes
MAN-DC-2-a		Outside		No	No	No	N <sub>o</sub>
MAN-UC-2-b	Dairy Cattle	Outside		No	No	No	ž
MAN-DC-3	1	Inside		No	Yes	No	ž
MAN-DC-3-b		Inside		No	Yes	No	%
MAN-DC-F	Field Blank	Outside		No	No	Νο	Š
MAN-SW-1-a	1	Outside		No	Yes	Yes	%
MAN-SW-I-b	Swine	Outside		No	Yes	No	Š
MAN-SW-2		Inside		No	No.	Νο	%

ioniai ner

STOK. 029772

Table 4. (continued).

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Did the Potential Biomarker Sequence PCR Amplify in the Fecal Sample? Clone SA15
E. coli Yes Yes Yes × Xes Yes Κes Kes Yes Yes ž ŝ Yes Yes å ŝ ş ž ž £ ŝ Ϋ́ ŝ ŝ å å ž Clone SA19 Kineococcus Yes, faint Yes Yes Yes Yes Yes Yes Xes Yes Yes ટ ટ્ટ ۶ 22 2 £ 원원 Yes X S ζeς. ž χes ટ્ટ å ž Unknown Organism Clone SB37 Yes Yes Yes Yes Yes Yes ž ş Yes Yes Yes Yes Yes Yes Kes Υes Yes ŝ å ဍ Yes Š Š. Yes Yes Νo Brevibacterium Clone LA35 Yes, fainthb No Yes Š Ŷ ž ž ž ž ટ g ž å å ပ္ပ ŝ ŝ ž ဥ No Š ž £ Š ž 22 Siloam Springs Siloam Springs Fayetteville Fayetteville Claremore Tulsa City Inside or Outside the Watershed a: Samples were re-extracted and re-run in duplicate to confirm result.
 b: Faint, indicates a very weak band was observed on the gel. Outside Outside Outside Outside Outside Outside Outside Outside Inside Outside Outside Outside Inside Inside Inside Inside Inside Inside Outside Inside Inside Inside Inside Inside Inside Inside Inside Type of Fecal Septic System Field Blank Waste Water Sample Treatment Goose Duck Plant MAN-DK-2-a MAN-DK-2-b MAN-DK-4-a MAN-DK-4-b MAN-DK-1-a MAN-DK-1-b MAN-DK-3-a MAN-DK-3-b MAN-DK-5-a MAN-DK-5-b MAN-GS-1-a MAN-GS-1-b MAN-GS-3-a MAN-GS-4-a MAN-GS-5-a MAN-GS-2-a MAN-GS-2-b MAN-GS-3-b MAN-GS-4-b MAN-GS-5-b MAN-DK-F MAN-HM-5 MAN-HM-2 MAN-HM-3 MAN-HM-4 MAN-HM-6 MAN-HM-I Sample

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### 3.4 Sequencing of the Duck and Goose Amplicon Derived from the LA35 Primer Set

#### 3.4.1 Objective

The purpose of the cloning and sequencing of the duck and goose amplicon derived from the LA35 primer set was to determine if these DNA sequences contained variable regions that could be targeted for a more specific set of PCR primers.

#### 3.4.2 **Method**

- Several clone libraries were constructed from the duck sample MAN-DK-1-b and the
  goose sample MAN-GS-1-a by PCR amplifying DNA from these fecal samples with the
  LA35 primer set, with primers 8F-907R and with 8F-1492R, and cloning with the
  TOPO-TA cloning kit (Invitrogen). The plasmids were excised from these clones
  (QIAprep Spin Miniprep Kit, QIAGEN).
- Extracted plasmids were DNA sequenced using the PCR primers T7, T3, 8F, 907R, or 1492R (as appropriate) by the MRCF. Sequences were compared to the poultry litter Brevibacteria biomarker using BioEdit V.7.0.5.3 to look for variable regions between the DNA sequences.

#### 3.4.3 Results

The regions of the 16S rRNA gene of the duck and goose DNA targeted by the poultry litter specific *Brevibacteria* biomarker PCR primers are identical to our biomarker *Brevibacteria* sequence.

After screening 20 clones from each of the two clone libraries developed (a total of 40 clones) from the duck and the goose fecal samples, only one organism was identified as a *Brevibacteria* using the BLAST database. This sequence was identical to the *Brevibacteria* biomarker sequence obtained from the original soil and litter samples within the 1,200 DNA base pairs sequenced.

## 3.5 Test for the Poultry Litter Specific Biomarker in Environmental Samples from Within the Impacted Watershed

#### 3.5.1 Objective

The purpose of this test was to determine if the *Brevibacteria* biomarker that is specific to poultry litter could be detected in environmental media (poultry litter, soil and water samples) from within a potentially poultry litter impacted watershed.

#### 3.5.2 Methods

Environmental samples were collected from within the potentially impacted watershed and
included chicken and turkey litter samples, soil samples from which the litter had been
applied, edge of field runoff water samples from the fields to which the litter had been
applied, river water to which the runoff samples drained, and lakes downgradient from the
previously collected river samples. Additionally, groundwater samples within the
potentially impacted watershed were collected and analyzed.

- Water samples were collected in duplicate 1-L sterile nalgene bottles and shipped on ice to the laboratory, where they were immediately filtered. The filters containing the microorganisms were frozen at -80°C until DNA extraction.
- Genomic DNA was extracted from the filters and soil and litter samples using the Bio 101 FastDNA SPIN Kit for Soil, DNA extraction kit (Qbiogene, Inc).
- Samples were tested for PCR amplification with the LA35 PCR primers, and products were separated by gel electrophoresis, stained with ethidium bromide, and visualized by UV light. Samples were run with a nested PCR approach

#### 3.5.3 Results

The results of the PCR amplification of the environmental samples for the poultry litter biomarker are presented in Table 5.

Table 5. Results of the PCR amplification of the environmental samples for the poultry litter biomarker.

Sample	Type of Sample	Amplified with LA35 PCR Primers?
FAC1-6-20-06	Litter	Yes
FAC2-6-21-06	Litter	Yes
FAC-03-7-6-06	Litter	Yes
FAC-4-7-12-06	Litter	Yes
FAC-5-7-13-06	Litter	Yes
FAC-6-7-20-06	Litter	No <sup>s</sup>
FAC-7-8-3-06	Litter	Yes
FAC-8-8-15-06	Litter	Yes
FAC-9-8-31-06	Litter	Yes
LAL5-A-2-6-13-06	Soil	No
LAL5-C-2-6-12-06	Soil	Yes
LAL-7-A-2-6-20-06	Soil	Yes
LAL-7-B-2-6-20-06	Soil	Yes, faint <sup>b</sup>
LAL-7-C-2-6-19-06	Soil	Yes, faint <sup>b</sup>
LAL10-B-2-6-26-06	Soil	No
LAL10-A-2-6-26-06	Soil	No
LAL10-A-4-6-26-06	Soil	No
LAL8-A-2-6-19-06	Soil	Yes
LAL8-B-2-6-21-06	Soil	No
LAL9-D-2-6-22-06	Soil	No
LAL9-B-2-6-22-06	Soil	Yes
LAL9-A-2-6-22-06	Soil	Yes
LAL7-D-2-6-29-06	Soil	Yes

Table 5. (	(continued)	).

Sample	Type of Sample	Amplified with Poultry Litter Specific Brevibacteria PCR Primers?
LAL8-D-2-6-20-06	Soil	Yes
LAL11-C-2-6-28-06	Soil	No
LAL11-D-2-6-28-06	Soil	No
LAL11-A-2-6-29-06	Soil	Yes
LAL11-D-2-Q-6-28-06	Soil	Yes
LAL12-A-2-7-6-06	Soil	No
LAL12-A-2-Q-7-6-06	Soil	Yes
LAL12-C-2-7-7-06	Soil	No
LAL12-D-2-7-7-06	Soil	No
LAL 13-A-2-7-6-06	Soil	Yes
LAL 13-C-2-7-7-06	Soil	No
LAL 13-C-2-Q-7-7-06	Soil	Yes
LAL 13-D-2-7-6-06	Soil	No
EOF-1-6-17-06	Water	No
EOF-Q2-6-17-06	Water	No
EOF-Q1-6-17-06	Water	Yes
EOF-SPREAD073B-6-18-06	Water	Yes
EOF-SPREAD023-6-18-06	Water	Yes
EOF-SPREAD044-6-18-06	Water	No
EOF-SPREAD068-6-18-06	Water	Yes

A summary of the presence or absence of the Brevibacteria biomarker in all samples analyzed to date is included in Table 6.

b: Faint indicates that a weak band was visible on the gel.

samples.

Type of Sample	Number Analyzed	Number Positive	% Containing LA35 sequence	Note
Original soil and litter samples	20	18	90.0	
Fecal Samples	57	2	3.5	LA35 sequences were present in one duck and one goose sample from outside the watershed
Additional litter samples	9	8	88.9	One of the litter samples (FAC-06) was taken from a poultry house without a concrete floor and likely contained a high percentage of soil.
Additional soil samples	27	14	51.9	
Edge of field water samples	7	4	57.1	

## 4 OPTIMIZATION AND VALIDATION OF A QUANTITIVE ASSAY FOR THE POULTRY LITTER SPECIFIC *BREVIBACTERIA* BIOMARKER

### 4.1 Develop a Quantitative PCR Assay for the *Brevibacterium* LA35 Biomarker

#### 4.1.1 Objective

The purpose of this work was to develop a qPCR assay for quantification of the LA35 biomarker in various environmental and fecal samples.

#### 4.1.2 Method

- LA35 PCR primers developed in Section 2 and validated for specificity to poultry litter in Section 3 were applied as qPCR primers using SYBR Green chemistry (dye) on a Chromo4 qPCR system (Bio-Rad).
- Plasmids containing the LA35 DNA sequence were used to create a standard curve, to
  determine a method detection limit (MDL) for positive control plasmids, and to determine
  the efficiency of the reaction. All standard curves were run in triplicate to verify the
  reproducibility of assay.
- Detection limits of the assay in environmental samples were determined by spiking soil
  and water samples with LA35 contained on plasmids. Controls were run to determine the
  average number of plasmids present in the E. coli cells used as plasmid carriers.
  Additionally, nanopure water and a composite water sample from the watershed were
  spiked with the E. coli containing the biomarker sequence and were filtered according to

the standard filtering methods. Cloning was performed using the Topo TA cloning kit (Invitrogen). Genomic DNA was extracted from the spiked samples using the Bio 101 FastDNA SPIN Kit for Soil, DNA extraction kit (Qbiogene, Inc). A graphic depicting the MDL protocol is presented in Figure 1.

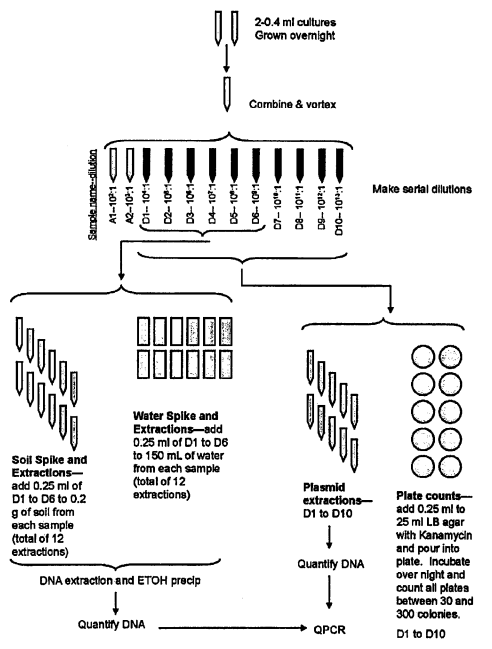
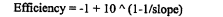


Figure 1. Outline of MDL protocol.

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#### 4.1.3 Results

The SYBR green qPCR standard curve is presented in Figure 2. Additionally, the plasmid MDL and reaction efficiency are presented in the figure. Efficiency of the qPCR reaction was determined by equation 1.





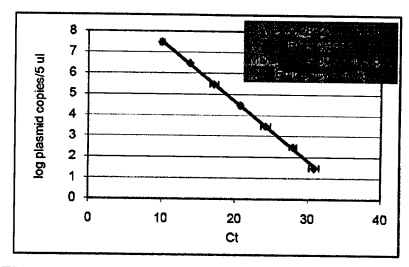


Figure 2. SYBR green qPCR standard curve of the poultry litter specific primers against plasmid DNA containing the *Brevibacteria* biomarker DNA. Error bars indicate the standard deviation of cycle thresholds of triplicate samples.

The results of the minimum detection limit test in spiked environmental samples are presented in Table 7.

Table 7. Detection limit of the qPCR assay for a poultry litter specific Brevibacteria in soils and water.

Sample Type	Minimum Detection Limit	Units
Plasmid DNA (standard)	6	copies/µL DNA extraction
Nanopure Water	18	cells/L
Composite Water Sample	78	cells/L
LAL11D-2Q-6-28-06 (soil) with sepharose cleanup	73	cells/g

### 4.2 Test qPCR Specificity to Distinguish Among Brevibacteria spp.

#### 4.2.1 Objective

The purpose of this test was to determine if the qPCR assay is specific enough to distinguish between the poultry litter biomarker and the closely related (but not identical) *Brevibacterium casei* 16S rRNA gene.

#### 4.2.2 Methods

A Brevibacterium casei culture was ordered from American Type Culture Collection (ATCC) and the DNA was extracted with the standard protocol. The extracted DNA was then tested for amplification with our qPCR protocol.

#### 4.2.3 Results

Our qPCR primers amplified the *Brevibacteria casei* 16S rRNA gene. As shown in Figure 3, we are able to distinguish *B. casei* from the LA35 sequenceby the SYBR green melt curves.

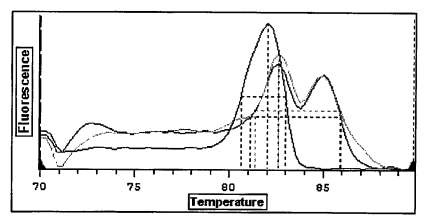


Figure 3. SYBR green melting curve profile of the LA35 16S rRNA sequence (blue) and that of *Brevibacterium casei* [ATCC 35513] (red and green).

## 4.3 Test for the Poultry Litter Specific Biomarker in Environmental Samples from Within the Impacted Watershed by the qPCR Assay

#### 4.3.1 Objective

The purpose of this test was to determine if the qPCR assay could detect the poultry litter specific biomarker in environmental samples from within the potentially impacted watershed.

#### 4.3.2 Method

- Genomic DNA was extracted from various environmental media (poultry litter, soil and water samples) per the previously described methods. Samples were selected that contained varying levels of fecal indicator bacteria, which was used to gauge the expected biomarker concentration (Table 8)
- DNA was subjected to a diagnostic PCR to verify that the samples did not contain contaminants that might inhibit qPCR and to determine the appropriate sample volume to use for the qPCR assay.
- DNA was then analyzed by the qPCR assay for the poultry litter specific biomarker. Any sample not amplifying in the qPCR assay was tested by a nested qPCR assay, where

#### 4.3.3 Results

The results of the qPCR and nested qPCR assay are presented in Table 8.

Sample ID	Matrix	Expected Biomarker Concentration	Expected DNA qPCR Poultry Specific Biomarker qPC Biomarker (ng/L or (copies/µL water or g soil or g Am Concentration ng/g)*	qPCR Poultry Specific Biomarker (copies/µL water or g soil or g litter)	Specifivater or litter)	CR Poultry Specific Biomarker (copies/µL water or g soll or g litter)	qPCR Matrix Spike Amplified?*	Nested qPCR Amplified?*	Biomarker Melt Peak Identified?	Other Meit Peaks Observed?
EOF-spr-010-5-9-06	Water	High	1.7	1.05E+07	₩	1.70E+06	Yes	N/A	Yes	οN
EOF-spr-17A-01-5-1-06	Water	High	72.5	2.48E+06	+1	4.71E+05	Yes	N/A	Yes	Yes
EOF-spr-023-6-18-06	Water	High	4.3	1.11E+05	++	2.49E+03	Yes	N/A	Yes	Š
EOF-spread-073B-6-18-06	Water	High	19.2	1.92E+06	#	4.42E+04	Yes	N/A	Yes	Š
LAL16-SPR2-7-18-06	Water	High	-1.0	Not present			Yes	No	N/A	N/A
LAL16C-2-7-18-06	Soil	High	9.5	1.42E+04	#	1.97E+03	Yes	N/A	Yes	No
LAL11C-2-6-28-06	Soil	High	73.2	Present, not quantifiable			Yes	Yes	Yes	Ν̈́ο
HFS16-BF1-01-6-15-06	Water	Medium	8.9	4.00E+03	#	1.60E+03	Yes	N/A	Yes	No
SALspr-6-28-06	Water	Medium	-0.6	5.82E+02	+1	1.56E+02	Yes	N/A	Yes	No
LAL15-SP2-7-11-06	Water	Medium	5.0	2.89E+03	#	7.69E+02	Yes	N/A	Yes	No
RS-PRICEcrk-01-4-29-06	Water	Medium	4.7	3.45E+05	#	1.43E+05	Yes	N/A	Yes	No
RS-574-BIO	Water	Medium	6.7	1.80E+05	#	6.09E+04	Yes	N/A	Yes	No
Lk04-0-01-5-16-06	Water	Low	8.9	3.69E+03	#	3.24E+03	Yes	N/A	Yes	No
HFS28A-BF1-01-6-15-06	Water	Low	-0.7	2.48E+03	#	1.28E+03	Yes	N/A	Yes	Yes
Rs-1-01-8-8-06	Water	Low	7.0	3.19E+04	#1	6.75E+03	Yes	N/A	Yes	Yes
FAC-01A-1	Litter	High	33.7	2.18E+09	#	3.53E+08	Yes	N/A	Yes	γ
FAC-01A-2	Litter	High	4.7	2.47E+08	#	3.22E+07	Υœ	N/A	Yes	%
FAC-01A-3	Litter	High	-0.5	2.67E+07	+1	2.69E+06	Yes	N/A	Yes	No

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Table 8. (continued).						-				
Sample ID	Matrix	Expected Biomarker Concentration	DNA (ng/L or ng/g)	qPCR Poultry Specific Biomarker (copies/µL water or g soil or g litter)	/ Specifi water oi litter)	ic Biomarker rg soil or g	qPCR Matrix Spike Amplified?*	Nested qPCR Amplified?*	Blomarker Melt Peak Identified?	Other Melt Peaks Observed?
FAC-01A-4	Litter	High	3.4	1.49E+08	-11	1.10E+07	Yes	N/A	Yes	No
FAC-01A-5	Litter	High	4.1	5.67E+08	#1	3.75E+07	Yes	N/A	Yes	No
FAC-01B-1	Litter	High	94.5	3.94E+09	#	6.28E+08	Yes	N/A	Yes	Ñ
FAC-01B-2	Litter	High	40.5	2.66E+09	#	7.57E+08	Yes	N/A	Yes	Ν̈́ο
FAC-01B-3	Litter	High	34.5	4.75E+06	+1	4.23E+06	Yes	N/A	Yes	No
FAC-01B-4	Litter	High	117.1	60+366'S	#	1.74E+09	Yes	N/A	Yes	SN.
LAL8-A-2-6-19-06	Soil	High	22.34	7.00E+03	+1	4.43E+02	Yes	N/A	Yes	No
LAL16B-2-7-18-06	Soil	High	28.94	2.91E+05	#1	1.95E+04	Yes	N/A	Yes	No
RS-901-BIO	Water	Low	1.3	Not present			Yes	Š	N/A	N/A
LAL16-GW2-7-18-06	Water	None	2.0	Not present			Yes	Š	N/A	N/A
CollinsWell#1-7-7-06	Water	None	4.0	Not present			səX	N <sub>O</sub>	N/A	N/A
66783-7-26-06	Water	None	8.0	Not present		•	Yes	Ν̈́	N/A	N/A
LK-01-0-01-8-9-06	Water	None	5.2	Not present			Yes	Š	N/A	N/A
Hester-498-8-10-06	Water	None	2.9	Not present			Yes	No	N/A	N/A
N/A, not applicable. The sample was not r	ple was not 1	run with the nested	qPCR assay	and/or the bioma	rker me	it peak was not	identified because	un with the nested qPCR assay and/or the biomarker melt peak was not identified because none was detected in the qPCR sample run.	in the qPCR samp	e run.
Inhibited indicates that the sample did not	mple did not		deven after a	ı sepharose clean	up was	performed and t	he sample was dil	amplify with qPCR even after a sepharose cleanup was performed and the sample was diluted to a lower DNA concentration.	A concentration.	
Concentration in water is given in ng/L; concentraiton in soil is in ng/g	ven in ng/L; (	concentraiton in soi	il is in ng/g							

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## 4.4 Determine the Effect of Dilution on Quantification of the LA35 Biomarker in Soil and Water Samples by the qPCR Assay

#### 4.4.1 Objective

This test was conducted to determine a concentration for poultry litter in an environmental sample below which the biomarker is likely not to be detected in environmental samples (i.e., simulate runoff conditions in the lab and test for the presence of biomarker at increasingly dilute samples).

#### 4.4.2 Methods

Litter samples FAC2 and FAC8 (see Table 5) were each carried through the entire procedure. Note that FAC2 was one of the original litter samples used to find LA35. Litter sample FAC8 is a turkey litter sample. Both FAC2 and FAC8 previously tested positive for the poultry litter specific biomarker by the qPCR assay. An outline of the protocol is presented in Figure 4.

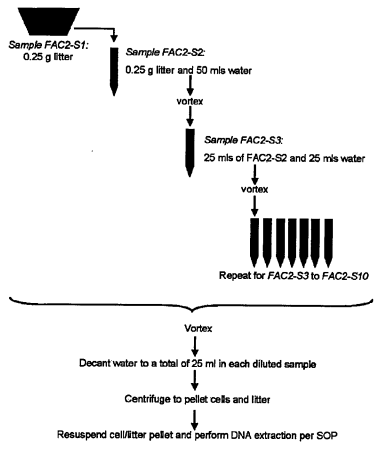


Figure 4. Outline of "dilution limit" protocol using litter sample FAC2 as the example.

#### 4.4.3 Results

LA35 was amplifiable in samples down to 0.1 mg of litter per L. The concentration of LA35 in the litter samples themselves was greater than  $2*10^8$  copies/g of litter. There was a very strong correlation between litter concentration (grams of litter per L) and the concentration of LA35 (copies of LA35 per gram of litter, as indicated by the  $R^2$  values of 0.97 and 0.99 shown in Figure 5. These  $R^2$  values indicate that the DNA extraction is efficient and *Brevibacteria* quantification method is relatively precise. Table 9 shows the results if the analysis.

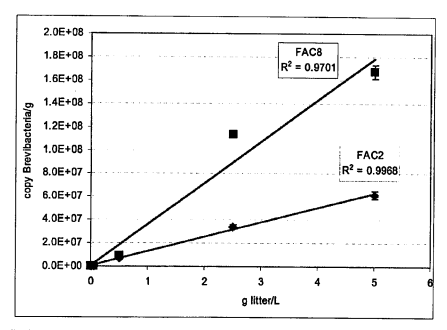


Figure 5. Copies of *Brevibacteria* poultry litter biomarker per gram of soil versus grams of soils per litter in the extractions.

Table 9. Results of the PCR, qPCR, and nested qPCR for Brevibacteria in litter.

			<del></del>			
Sample	"Litter" Concentration^	PCR Bacteria*	Nested qPCR*	Brevibacte		
FAC2-S1	NA	+	NA	2.32E+08	±	8.88E+06
FAC2-S2	5,000 mg/L	+	NA	6.14E+07	±	3.37E+06
FAC2-S3	2,500 mg/L	+	NA	3.42E+07	±	1.61E+06
FAC2-S4	500 mg/L	+	NA	6.86E+06	±	8.27E+05
FAC2-S5	50 mg/L	-	NA	3.54E+04	±	4.84E+03
FAC2-S6	10 mg/L	-	NA	1.90E+04	Ŧ	4.79E+03
FAC2-S7	5 mg/L		NA	1.01E+04	±	9.35E+03
FAC2-S8	1 mg/L	-	NA	Present, no	ot qu	antifiable

Table 9. (continued).

Sample	"Litter" Concentration^	PCR Bacteria*	Nested qPCR*	Brevibacteria 16S rRNA (copy/L water or g litter)
FAC2-S9	0.1 mg/L	-	+	Present, not quantifiable
FAC2-S10	0.01 mg/L	•	-	Not detected
FAC8-S1	NA	+	NA	2.56E+08 ± 2.49E+07
FAC8-S2	5,000 mg/L	+	NA	1.67E+08 ± 5.88E+06
FAC8-S3	2,500 mg/L	+	NA	1.14E+08 ± 1.56E+05
FAC8-S4	500 mg/L	+	NA	9.05E+06 ± 8.69E+05
FAC8-S5	50 mg/L	-	NA	7.48E+03 ± 2.53E+03
FAC8-S6	10 mg/L	-	NA	7.59E+04 ± 2.97E+04
FAC8-S7	5 mg/L	-	NA	3.20E+04 ± 2.23E+04
FAC8-S8	1 mg/L	-	+	Present, not quantifiable
FAC8-S9	0.1 mg/L	-	-	Not detected
FAC8-S10	0.01 mg/L	-	-	Not detected

<sup>^</sup> NA indicates that this was the original 0.25 g undiluted litter sample from which the dilutions were made.

#### 5 REFERENCES

Bernhard, A.E. and K.G. Field, 2000, "Identification of nonpoint sources of fecal pollution in coastal waters by using host-specific 16S ribosomal DNA genetic markers from fecal anaerobes," *Applied and Environmental Microbiology.* 66(4): p. 1,587-1,594.

Tsen, H, et al., 1998, "Development and use of 16S rRNA gene targeted PCR primers for the identification of Escherichia coli cells in water," *Journal of Applied Microbiology*. **85**(3): p. 554-560.

<sup># &</sup>quot;+" indicates that PCR products were observed after gel electrophoresis, indicating that amplifiable products were obtained from the sample. "-" indicates that PCR products were not observed by gel electrophoresis and UV visualization. Note that the detection limits of the gel electrophoresis method are much higher than the qPCR method detection limits.

<sup>\* &</sup>quot;4" indicates that the Brevibacteria were detected with the nested qPCR protocol. "NA" indicates that the samples were not tested with the nested qPCR protocol.



### Poultry-Specific Biomarker Quantitative PCR Analytical Summary

November 3, 2007

#### Overview:

The objective of this project was to quantify the number of poultry-specific *Brevibacteria* biomarker gene copies contained in water, soil, and/or litter samples using quantitative polymerase chain reaction (qPCR). Table 1 describes the sample matrix and the condition of the samples upon arrival to the analytical laboratory.

Table 1. Description of samples and volume or mass filtered for DNA extraction.

Sample ID	Matrix/ Date Sampled	Condition Received/Observations	Volume Filtered (mL) or Mass Extracted (g)
FAC03-7-6-06	Litter/7-6-06	Cold/sealed bag	0.35 g
FAC09-8-31-06	Litter/8-31-06	Cold/sealed bag	0.22 g
FAC02-6-21-06	Litter/6-21-06	Cold/sealed bag	0.24 g
FAC08-8-15-06	Litter/8-15-06	Cold/sealed bag	0.26 g
FAC04-7-12-06	Litter/7-12-06	Cold/sealed bag	0.26 g
FAC010-9-22-06	Litter/9-22-06	Cold/sealed bag	0.2 g
FAC07-8-3-06	Litter/8-3-06	Cold/sealed bag	0.39 g
FAC05-7-13-06	Litter/7-13-06	Cold/sealed bag	0.35 g
FAC06-7-20-06	Litter/7-20-06	Cold/sealed bag	0.27 g
LAL6A2-6-14-06	Soil/6-14-06	Cold/sealed bag	0.24 g
LAL6D2-6-15-06	Soil/6-15-06	Cold/sealed bag	0.24 g
LAL6B2-6-14-06	Soil/6-14-06	Cold/sealed bag	0.23 g
LAL13C2Q-7-7-06	Soil/7-7-06	Cold/sealed bag	0.24 g
LAL14C2-7-10-06	Soil/7-10-06	Cold/sealed bag	0.58 g
LAL14C2Q-7-10-06	Soil/7-10-06	Cold/sealed bag	0.51 g
LAL13C2-7-7-06	Soil/7-7-06	Cold/sealed bag	0.35 g
LAL16C2-7-18-06	Soil/7-18-06	Cold/sealed bag	0.35 g
LAL5C2-6-12-06	Soil/6-12-06	Cold/sealed bag	0.25 g
LAL8B2-6-21-06	Soil/6-21-06	Cold/sealed bag	0.26 g
LAL14B2-7-10-06	Soil/7-10-06	Cold/sealed bag	0.58 g
LAL9D2-6-22-06	Soil/6-22-06	Cold/sealed bag	0.25 g
LAL7B2-6-20-06	Soil/6-20-06	Cold/sealed bag	0.25 g
LAL13A2-7-6-06	Soil/7-6-06	Cold/sealed bag	0.27 g
LAL17C2-7-18-06	Soil/7-18-06	Cold/sealed bag	0.71 g
LAL17C2Q-7-18-06	Soil/7-18-06	Cold/sealed bag	0.62 g
LAL14D2-7-10-06	Soil/7-10-06	Cold/sealed bag	0.63 g
LAL8A2-6-19-06	Soil/6-19-06	Cold/sealed bag	0.7 g



orania de la compansión	B/s-4/	Candition	Volume Filtered (mT) on
Sample ID	Matrix/ Date Sampled	Condition Received/Observations	Volume Filtered (mL) or Mass Extracted (g)
LAL8D2-6-20-06	Soil/6-20-06	Cold/sealed bag	0.26 g
LAL17A2-7-10-06	Soil/7-10-06	Cold/sealed bag	0.68 g
LAL7D2-6-29-06	Soil/6-29-06	Cold/sealed bag	0.24 g
LAL9A2-6-22-06	Soil/6-22-06	Cold/sealed bag	0.26 g
LAL5A2-6-13-06	Soil/6-13-06	Cold/sealed bag	0.35 g
LAL7C2-6-19-06	Soil/6-19-06	Cold/sealed bag	0.25 g
LAL9B2-6-22-06	Soil/6-22-06	Cold/sealed bag	0.25 g
LAL13D2-7-6-06	Soil/7-6-06	Cold/sealed bag	0.24 g
LAL7A2-6-20-06	Soil/6-20-06	Cold/sealed bag	0.24 g
LAL16D2-7-18-06	Soil/7-18-06	Cold/sealed bag	0.52 g
LAL5B2-6-12-06	Soil/6-12-06	Cold/sealed bag	0.25 g
LAL12A2Q-7-6-06	Soil/7-6-06	Cold/sealed bag	0.27 g
LAL12D2-7-7-06	Soil/7-7-06	Cold/sealed bag	0.24 g
LAL16-SP2-7-18-06	Water/7-18-06	Cold/bottle intact	100 mL
EOF-SPREAD-010-5-9-06	Water/5-9-06	Cold/bottle intact	40 mL
EOF-SPREAD-17A-01-5-1-06	Water/5-1-06	Cold/bottle intact	30 mL
EOF-SPREAD-023-6-18-06	Water/6-18-06	Cold/bottle intact	25 mL
EOF-SPREAD-073B-6-18-06	Water/6-18-06	Cold/bottle intact	10 mL
EOF-SPREAD-064-5-4-06	Water/5-4-06	Cold/bottle intact	50 mL
EOF-SPREAD-53E-01-4-29-06	Water/4-29-06	Cold/bottle intact	30 mL
EOF-SPREAD-60-01-4-29-06	Water/4-29-06	Cold/bottle intact	50 mL
SPREAD-023-4-25-06	Water/4-25-06	Cold/bottle intact	40 mL
EOF-1-6-17-06	Water/6-17-06	Cold/bottle intact	100 mL
EOF-SPREAD-053G-5-4-06	Water/5-4-06	Cold/bottle intact	100 mL
EOF-SPREAD-048-5-9-06	Water/5-9-06	Cold/bottle intact	100 mL
SPREAD-029-4-25-06	Water/4-25-06	Cold/bottle intact	100 mL
SPREAD-036-4-25-06	Water/4-25-06	Cold/bottle intact	100 mL
EOF-SPREAD-071-5-9-06	Water/5-9-06	Cold/bottle intact	150 mL
EOF-SPREAD-065-5-4-06	Water/5-4-06	Cold/bottle intact	100 mL
EOF-Q2-6-17-06	Water/6-17-06	Cold/bottle intact	50 mL
EOF-26-6-8-05	Water/6-8-05	Cold/bottle intact	500 ml
EOF-17-6-8-05	Water/6-8-05	Cold/bottle intact	500 mL
EOF-222-4-13-07	Water/4-13-07	Cold/bottle intact	20 mL
GPGW-18A-6-26-07	Water/6-26-07	Cold/bottle intact	20 mL
GPGW-20-6-11-30-06	Water/11-30-06	Cold/bottle intact	50 mL
GPGW-48-7-12-1-06	Water/12-1-06	Cold/bottle intact	50 mL
GPGW-10-4-11-30-06	Water/11-30-06	Cold/bottle intact	50 mL





Sample ID	Matrix/ Date Sampled	Condition Received/Observations	Volume Filtered (mL) or Mass Extracted (g)
GPGW-40-6-27-07	Water/6-27-07	Cold/bottle intact	15 mL
HFS22-EVENTA-5-10-06	Water/5-10-06	Cold/bottle intact	150 mL
HFS04-BF2-01-8-1-06	Water/8-1-06	Cold/bottle intact	500 mL



#### Methods:

DNA Extraction. For soil and/or litter samples, DNA was extracted from 0.25 g of soil or litter using the FastDNA®SPIN® Kit for soil protocol. For surface water shipped to the laboratory, between 100 and 1,000 mL of groundwater was filtered through a Supor-200, 0.2 µm filter. The filters were frozen at -80°C and then shattered. Next, each sample tube was amended with 2 mL of DNA-free water, vortexed vigorously for 15 minutes, and the liquid volume was partitioned into DNA extraction tubes. DNA extractions were performed using the FastDNA®SPIN® Kit for soil according to the manufacturer's instructions. All DNA extractions were cleaned using an ethanol precipitation method. Community DNA was eluted in nucleasefree water (50 µL) and stored at -20°C.

Amplification of Bacteria. The PCR was used to amplify nearly full-length 16S rDNA genes from Bacteria. Each 25-μL PCR reaction included 1 X PCR buffer, 1.5 mM MgCl<sub>2</sub>, 0.5 μM each 8F forward and 907R reverse primer, 1 u/50uL Taq DNA polymerase, 0.2 mM dNTP, 1 uL template DNA, and 20 uL molecular-grade water. Amplification was performed on a MJ Research Peltier Gradient thermocycler using the following regime: 94°C (5 min) followed by 30 cycles of 94°C (1 min), 53.5°C (1 min), and 72°C (1 min, 50 sec). The reaction was finished with an additional 7 minutes at 72°C. PCR products were examined by ultraviolet (UV) light in a 1% agarose gel stained with ethidium bromide to confirm specificity of the amplification reactions.

Sepharose Cleanup. Any sample not amplifying in the PCR was processed through a Sepharose CL-4B (Sigma-Aldrich) size exclusion gel chromatography cleanup. Briefly, the micro-bio spin columns (Bio-Rad) were packed with sterile Sepharose CL-4B and washed with Tris-HCl buffer (pH 8). The sample was added to the packed gel column and eluted by spinning in a micro-centrifuge.

Detection of a Poultry Specific Brevibacteria Biomarker. The qPCR methods for assessing the 16S rRNA gene are very sensitive in detecting specific DNA fragments. The detection limit for the methods used is approximately 6 gene copies per µL of the DNA extraction. Biomarker DNA was cloned into a plasmid and was used as the source of the quantitative standards used in the analysis. Plasmid DNA containing the target 16S rRNA gene from the poultry-specific Brevibacteria biomarker was purified and quantified fluorometrically. Based on the known size of the plasmid and insert, DNA concentrations were converted to insert copy numbers. A dilution series spanning seven orders of magnitude was generated using known concentrations of each plasmid. Amplification and detection of the DNA was performed using the MI Chromo-4 System. The acceptance criterion for the standard curve is a linear R<sup>2</sup> value of greater than 0.995.

To determine qPCR results, sample DNA diluted to a final concentration of 15 ng/5 μL DNA was combined with following reagents to reach a final concentration of 1X SYBR Green Master Mix and 0.5 µM 157F and 727R primer and water to reach 20  $\mu L$  and 5  $\mu L$ , respectively, of diluted sample DNA. Amplification was performed on the MJ Research PTC-2004 thermocylcer using the following regime: 50°C (2 min), 95°C (15 min), 40 cycles of 95°C (30 sec), 60°C (1 min), plate read and 50°C (5 min). The melting curve was determined using the following protocol: heat from 60°C to 90°C, by 0.3°C increments, holding for 5 seconds before reading the fluorescence of the samples. Nested qPCR results were determined by purifying the PCR products using the QIAquick PCR Purification Kit, as per the manufacturer's protocol, and then running the purified samples through qPCR, as described above.

QA/QC Requirements. To determine if and where potential contamination or interference occurred during sample processing, positives and reagent blanks or negatives and matrix spikes of the PCR and qPCR samples were prepared. A positive control consisting of pure DNA (known to amplify by specific DNA primers) was used for the PCR and qPCR procedure. A matrix spike consisting of pure DNA (known to amplify by specific DNA primers) was used for the PCR and qPCR procedure. Negative controls consisted of water-only blanks for the PCR and qPCR procedure. The qPCR reactions were run in triplicate for each sample to determine the reproducibility of the method.



# Results:

subjected to polymerase chain reaction (PCR) using universal bacterial probes in order to verify amplifiable DNA was present in the sample. In addition, The samples arrived at the lab in good condition at 4°C with ice still in the cooler. The samples were filtered in the lab, and the filters were for the 16S rRNA gene, a "nested" qPCR approach was applied in which the universal bacterial PCR-amplified DNA is used as the template in a qPCR reaction. Although the results from the nested qPCR cannot be quantified per se, they can be used to lower the detect limit for the qPCR in order to groundwater DNA extractions. The results of these studies are presented in Table 2. The DNA extraction negative control and all PCR negative immediately placed in a -80°C freezer and stored until the DNA extraction was performed. Following DNA extraction, the samples were first determine if the poultry-specific Brevibacteria biomarker gene is present at concentrations lower than the method detect limit (MDL) using the controls did not amplify any product. In addition, all calibration control checks were within acceptable values.

inhibition was removed in the second cleanup and the Brevibacteria biomarker was detected in the qPCR analysis. The Brevibacteria biomarker reported in September. This sample was sepharose cleaned an additional time and reanalyzed per the standard operating procedure (SOP). All Sample LAL16C2-7-18-06 results were reported previously (September 17, 2007). This sample was inhibited in the prior analysis as was identified in 81% of the samples and was quantifiable in 39% of the samples analyzed.

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Matrix n		qPCR Poultry Specific Biomarker (copies/µL), water soil or o litter)*	ry Spe	ecific Frater or g	qPCR Matrix Spike	Nested qPCR	Biomarker Melt Peak	Other M. Peaks
Litter	21.3	1.03E+09	#1	8.00E+07	Yes	N/A	Yes	S <sub>N</sub>
Litter 1	70.1	7.57E+08	#	1.55E+08	Yes	N/A	Yes	å
Litter :	51.9	4.13E+08	#	1.78E+07	Yes	N/A	Yes	%
Litter 1	54.0	1.47E+09	#	2.25E+08	Yes	N/A	Yes	%
Litter	8.9	1.67E+08	#	2.98E+07	Yes	N/A	Yes	%
Litter 1	20.1	2.04E+09	+1	4.14E+08	Yes	N/A	Yes	%
Litter 5	98.1	2.49E+09	#	9.54E+07	Yes	N/A	Yes	%
Litter 7	9.9/	1.47E+09	#	1.93E+08	Yes	N/A	Yes	Š
Litter	57.1	4.46E+08	+1	7.34E+07	Yes	N/A	Yes	% N
Soil	10.5	1.55E+04	#	2.57E+03	Yes	N/A	Yes	%
- +:e:e:e:e:e:e:e:e:e:e:e:		(ng/L or ng/g) 21.3 21.3 170.1 51.9 154.0 6.8 6.8 120.1 98.1 76.6 57.1	(ng/L or ng/g) 21.3 21.3 170.1 51.9 154.0 6.8 6.8 120.1 98.1 76.6 57.1	(ng/L or ng/g) 21.3 21.3 170.1 51.9 154.0 6.8 6.8 120.1 98.1 76.6 57.1	(ng/L or Biomarker (coples/µL w ng/g)  21.3 1.03E+09 ± 8  170.1 7.57E+08 ± 8  51.9 4.13E+09 ± 8  6.8 1.67E+09 ± 8  120.1 2.04E+09 ± 8  76.6 1.47E+09 ± 6  76.6 1.47E+09 ± 6  76.6 1.47E+09 ± 7  76.6 1.45E+09 ± 6  76.6 1.45E+09 ± 7  76.6 1.45E+	(ng/L or ng/g)         Biomarker (coples/µL) water or g soil or g litter)*           21.3         1.03E+09         ±         8.00E+07           170.1         7.57E+08         ±         1.55E+08           51.9         4.13E+08         ±         1.78E+07           154.0         1.47E+09         ±         2.25E+08           6.8         1.67E+08         ±         2.98E+07           120.1         2.04E+09         ±         4.14E+08           98.1         2.49E+09         ±         9.54E+07           76.6         1.47E+09         ±         1.93E+08           57.1         4.46E+08         ±         7.34E+07           10.5         1.55E+04         ±         2.57E+03	(ng/L or ng/g)         Biomarker (coples/µL water or g soil or g litter)*         Splike Amplified?*           21.3         1.03E+09         ±         8.00E+07         Yes           170.1         7.57E+08         ±         1.55E+08         Yes           51.9         4.13E+09         ±         1.75E+08         Yes           154.0         1.47E+09         ±         2.25E+08         Yes           6.8         1.67E+08         ±         2.98E+07         Yes           120.1         2.04E+09         ±         4.14E+08         Yes           98.1         2.49E+09         ±         9.54E+07         Yes           76.6         1.47E+09         ±         1.93E+08         Yes           57.1         4.46E+08         ±         7.34E+07         Yes           10.5         1.55E+04         ±         2.57E+03         Yes	(ng/L or ng/L)         Biomarker (coples/µL) water or g soil or g litter)*         Spike Amplified?*         N/A         N/A

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		DNA (ng/L or	qPCR Poultry Specific Biomarker (copies/µL water or g	try Sp es/µL	ecific water or g	qPCR Matrix Spike	Nested aPCR	Biomarker Melt Peak	Other Melt Peaks
Sample ID	Matrix	ng/g)	soil or g litter)	litter	,	Amplified? <sup>b</sup>	Amplified?	Identified?	Observed?
LAL6D2-6-15-06	Soil	2.1	4.98E+03	#1	1.88E+02	Yes	N/A	Yes	No
LAL6B2-6-14-06	Soil	12.4	Present			Yes	Yes	Yes	ž
LAL13C2Q-7-7-06	Soil	27.3	Present			Yes	Yes	Yes	ž
LAL14C2-7-10-06	Soil	20.3	Present			Yes	Yes	Yes	%
LAL14C2Q-7-10-06	Soil	77.3	Present			Yes	Yes	Yes	8
LAL13C2-7-7-06	Soil	3.4	Present			Yes	Yes	Yes	%
LAL16C2-7-18-06	Soil	5.6	1.42E+04	#	1.97E+03	Yes	N/A	Yes	ν
LAL5C2-6-12-06	Soil	40.3	Present			Yes	Yes	Yes	%
LAL8B2-6-21-06	Soil	27.7	0.00E+00			Yes	No	N/A	N/A
LAL14B2-7-10-06	Soil	91.7	Present			Yes	Yes	Yes	ž
LAL9D2-6-22-06	Soil	13.0	Present			Yes	Yes	Yes	Š
LAL7B2-6-20-06	Soil	47.4	Present			Yes	Yes	Yes	No
LAL13A2-7-6-06	Soil	38.6	Present			Yes	Yes	Yes	No
LAL17C2-7-18-06	Soil	17.3	Present			Yes	Yes	Yes	No
LAL17C2Q-7-18-06	Soil	29.9	Present			Yes	Yes	Yes	No
LAL14D2-7-10-06	Soil	28.1	Present			Yes	Yes	Yes	No
LAL8A2-6-19-06	Soil	22.3	7.00E+03	#	4.43E+02	Yes	N/A	Yes	S.
LAL8D2-6-20-06	Soil	38.4	0.00E+00			Yes	No	N/A	N/A
LAL17A2-7-10-06	Soil	21.4	Present			Yes	Yes	Yes	%
LAL7D2-6-29-06	Soil	20.6	Present			Yes	Yes	Yes	% S
LAL9A2-6-22-06	Soil	29.2	Present			Yes	Yes	Yes	No
LAL5A2-6-13-06	Soil	35.9	Present			Yes	Yes	Yes	Š
LAL7C2-6-19-06	Soil	41.8	Present			Yes	Yes	Yes	No
LAL9B2-6-22-06	Soil	61.4	Present			Yes	Yes	Yes	No
LAL13D2-7-6-06	Soil	34.6	Present			Yes	Yes	Yes	No No



		DNA	aPCR Poultry Specific	try Sp	ecific	oPCR Matrix		Biomarker	Other Melt
Sample ID	Matrix	(ng/L or ng/g)	Biomarker (copies/µL water or soil or g litter)*	es/µL	water or g	Spike Amplified? <sup>b</sup>	Nested qPCR Amplified?	Melt Peak Identified?	Peaks Observed?
LAL7A2-6-20-06	Soil	31.7	Present			Yes	Yes	Yes	Νο
LAL16D2-7-18-06	Soil	36.4	Present			Yes	Yes	Yes	No
LAL5B2-6-12-06	Soil	27.3	Present			Yes	Yes	Yes	No
LAL12A2Q-7-6-06	Soil	26.8	Present			Yes	Yes	Yes	No
LAL12D2-7-7-06	Soil	51.4	Present			Yes	Yes	Yes	Ño
LAL16-SP2-7-18-06	Water	-1.0	0.00E+00			Yes	ν	N/A	N/A
EOF-SPREAD-010-5-9-06	Water	1.7	1.05E+07	H	1.70E+06	Yes	N/A	Yes	No
EOF-SPREAD-17A-01-5-1-06	Water	72.5	2.48E+06	#	4.71E+05	Yes	N/A	Yes	Yes
EOF-SPREAD-023-6-18-06	Water	4.3	1.11E+05	Ħ	2.49E+03	Yes	N/A	Yes	No
EOF-SPREAD-073B-6-18-06	Water	133.5	5.56E+07	#1	5.25E+06	Yes	N/A	Yes	S <sub>S</sub>
EOF-SPREAD-064-5-4-06	Water	79.2	1.89E+06	#1	7.63E+04	Yes	N/A	Yes	No
EOF-SPREAD-53E-01-4-29-06	Water	57.7	5.45E+07	H	4.80E+06	Yes	N/A	Yes	δ
EOF-SPREAD-60-01-4-29-06	Water	431.4	3.90E+07	#	8.26E+06	Yes	N/A	Yes	No
SPREAD-023-4-25-06	Water	194.2	1.25E+06	#	2.35E+05	Yes	N/A	Yes	No
EOF-1-6-17-06	Water	2.5	1.15E+05	#	1.80E+04	Yes	N/A	Yes	No
EOF-SPREAD-053G-5-4-06	Water	14.5	0.00E+00			Yes	No	N/A	N/A
EOF-SPREAD-048-5-9-06	Water	25.2	0.00E+00			Yes	No	N/A	N/A
SPREAD-029-4-25-06	Water	56.1	Present			Yes	Yes	Yes	No
SPREAD-036-4-25-06	Water	64.9	1.48E+05	#	4.04E+04	Yes	N/A	Yes	οN
EOF-SPREAD-071-5-9-06	Water	5.2	3.63E+04	#	8.25E+03	Yes	N/A	Yes	No
EOF-SPREAD-065-5-4-06	Water	13.3	3.45E+04	#	1.64E+03	Yes	N/A	Yes	No
EOF-Q2-6-17-06	Water	2.0	0.00E+00			Yes	Ño	N/A	N/A
EOF-26-6-8-05	Water	14.3	0.00E+00			Yes	No	N/A	N/A
EOF-17-6-8-05	Water	22.7	Present			Yes	Yes	Yes	No
EOF-222-4-13-07	Water	81.3	1.32E+05	Ŧ	2.71E+04	Yes	N/A	Yes	No
GPGW-18A-6-26-07	Water	1.1	0.00E+00			Yes	No	N/A	N/A



		DNA (ng/L or	qPCR Poultry Specific Biomarker (copies/µL water or g	/ Specific	qPCR Matrix Spike	Nested qPCR	Biomarker Meit Peak	Other Melt Peaks
Sample ID	Matrix	ng/g)	soil or g litter)*	tter)*	Amplified? <sup>b</sup>	Amplified?	Identified?	Observed?
GPGW-20-6-11-30-06	Water	13.4	0.00E+00		Xes.	oN	N/A	N/A
GPGW-48-7-12-1-06	Water	2.6	0.00E+00		Yes	No	N/A	N/A
GPGW-10-4-11-30-06	Water	2.9	Present		səX	Yes	Yes	SN.
GPGW-40-6-27-07	Water	-1.5	0.00E+00		səA	No	N/A	N/A
HFS22-EVENTA-5-10-06	Water	4.2	0.00E+00		Хcs	No	N/A	N/A
HFS04-BF2-01-8-1-06	Water	0.0	0.00E+00		Yes	No	N/A	N/A
• "Present" indicates that the biomarker was amplified, but was not quantifiable.  bif "no" indicates that sample did not amplify with qPCR even after a sepharose clearup was performed and the sample was diluted to a lower DNA concentration indicative of inhibition.  N/A, not applicable. The sample was not run with the nested qPCR assay and/or the biomarker melt peak was not identified because none was detected in the qPCR sample run.	was amplified, t mplify with qPCI not run with the r	amplified, but was not quantifiable.  fy with qPCR even after a sepharose un with the nested qPCR assay and/	amplified, but was not quantifiable. If with qPCR even after a sepharose cleamp was performed and the sample was diluted to a lower DNA concentration indicative of it un with the nested qPCR assay and/or the biomarker melt peak was not identified because none was detected in the qPCR sample run.	rformed and the san melt peak was not	nple was diluted to a lov identified because none	ver DNA concentratio was detected in the qI	n indicative of inhibi PCR sample run.	tion.